ACCESSION NUMBER:

1998:130639 CAPLUS Full-text

DOCUMENT NUMBER:

128:228387

TITLE:

Differential effects of aminosubstituted analogs of

hydroxy bisphosphonates on the growth of Dictyostelium

discoideum

AUTHOR (S):

Brown, R. J.; Van Beek, E.; Watts, D. J.; Lowik, C. W.

G. M.; Papapoulos, S. E.

CORPORATE SOURCE:

Department of Molecular Biology and Biotechnology,

University of Sheffield, Sheffield, UK

SOURCE:

Journal of Bone and Mineral Research (1998), 13(2),

253-258

CODEN: JBMREJ; ISSN: 0884-0431

PUBLISHER:

Blackwell Science, Inc.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

AB Replacing the hydroxyl group in the bone-binding site of three clin. useful bisphosphonates (etidronate, pamidronate, and olpadronate) by an amino group resulted in great differences in their antiresorptive potencies in vitro. the present study, this is also shown in vivo in mice treated with the six bisphosphonates at doses of up to 16 µM/kg/day for 12 days. Because binding to bone mineral is nearly the same for all tested bisphosphonates, these findings suggest that the aminosubstitution affects the cellular action of the bisphosphonates. This was tested in the cellular slime mold Dictyostelium discoideum in which cellular effects of bisphosphonates can be examined independently of binding to bone mineral. Etidronate and its aminosubstituted analog were equipotent in inhibiting amebal growth, while pamidronate was somewhat more potent than its analog. Whereas olpadronate was a potent inhibitor of axenic growth of Dictyostelium amebae, the aminosubstitution reduced its potency drastically (IC50 12 μM and 700 μM , resp.). similarities between the inhibitory effects of the bisphosphonates tested on bone resorption in vitro and in vivo and on the growth of Dictyostelium amebae confirm that the differences in antiresorptive potencies found reflect differences in cellular effects and suggest that bisphosphonates may bind to more than one intracellular target.

IT 63132-38-7

> RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)

(differential effects of amino-substituted analogs of hydroxy bisphosphonates on growth of Dictyostelium discoideum)

RN 63132-38-7 CAPLUS

CN Phosphonic acid, P,P'-[1-amino-3-(dimethylamino)propylidene]bis-INDEX NAME)

REFERENCE COUNT:

THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS 14 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L38 ANSWER 9 OF 11 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1997:132766 CAPLUS Full-text

DOCUMENT NUMBER:

126:144414

TITLE: INVENTOR(S): Amino-substituted bisphosphonic acids Papapoulos, Socrates; Van Beek, E. R.; Lowick, C. W. G. M.; Labriola, Rafael; Vecchioli,

Adriana

PATENT ASSIGNEE(S):

Gador S.A., Argent.; University of Leiden

SOURCE:

Eur. Pat. Appl., 14 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| | PA: | ENT | NO. | * | | KIN | D | DATE | | | APP | LICAT | 'ION | NO. | | D. | ATE | |
|-------|-----|-------|------|------------|-----|---------|-----|------|------|-----|-----|-------|------|-----|------|-----|------|-----|
| | | | | - - | | | - | | | | | | | | | - | | |
| | EΡ | 7535 | 23 | | | . A1 | | 1997 | 0115 | | EP | 1995- | 1107 | 06 | | 1 | 9950 | 710 |
| | | R: | GB | | | | | | | | | | | | | | | |
| | WO | 9702 | 827 | | | A1 | | 1997 | 0130 | | WO | 1996- | EP29 | 81 | | 1 | 9960 | 708 |
| | | W: | AU, | BR, | CA, | CN, | CZ, | FI, | IL, | JP, | ΚP | , KR, | NO, | PL, | -RU, | SK, | US, | VN |
| | | RW: | AT, | BE, | CH, | DE, | DK, | ES, | FI, | FR, | GB | , GR, | ΙE, | IT, | LU, | MC, | NL, | PT, |
| | | | SE, | BF, | ВJ, | CF, | CG, | CI, | CM, | GΑ, | GN | , ML, | MR, | NE, | SN, | TD, | TG. | |
| | ΑU | 9666 | 125 | | | Α | | 1997 | 0210 | | AU | 1996- | 6612 | 5 | | 1 | 9960 | 708 |
| | ΕP | 8376 | 82 | | | A1 | | 1998 | 0429 | | ΕP | 1996- | 9256 | 79 | | 1 | 9960 | 708 |
| | ΕP | 8376 | 82 | | | B1 | | 2002 | 1106 | | | | | | | | | |
| | | R: | DE, | FR, | GB, | NL | | | | | | | | | | | | |
| | JΡ | 1150 | 8905 | | | ${f T}$ | | 1999 | 0803 | | JP | 1996- | 5054 | 94 | | 1 | 9960 | 708 |
| | ZA | 9605 | 798 | | | Α | | 1998 | 0109 | | zA | 1996- | 5798 | | | 1 | 9960 | 709 |
| | US | .5990 | 098 | | | Α | | 1999 | 1123 | | US | 1998- | 9832 | 47 | | 1 | 9980 | 901 |
| PRIOR | IT | APP | LN. | INFO | .: | | | | | | ΕP | 1995- | 1107 | 06 | 2 | A 1 | 9950 | 710 |
| | | | | | | | | | | | WO | 1996- | EP29 | 81 | 1 | W 1 | 9960 | 708 |

OTHER SOURCE(S): MARPAT 126:144414

AB 1-Aminoalkylidene-1,1-bisphosphonic acids, RC(NH2)[P(O)(OH)2]2 (R = C1-9 straight-chain or branched aliphatic hydrocarbon radical which is optionally substituted by one or more amino or aminoalkyl groups with the exception of a terminal aminoalkyl group NR1R2; R1 = C1-9 straight-chain or branched, saturated or unsatd. aliphatic hydrocarbon radical, R2 = cyclohexyl or cyclohexylmethyl, benzyl or a straight-chain or branched, C4-18 saturated or unsatd. aliphatic hydrocarbon radical, as a single substituent of R) or any salts thereof, useful for treatment of disorders of calcium and bone metabolism, is described. Thus, hydrolysis of PC13 gave phosphorus acid which on treatment with MeCN in MeOH followed by acidic workup gave 100% MeC(NH2)[P(O)(OH)2]2. Some binding of compds. prepared with bone materials is described.

IT 63132-38-7P

> RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation and bone binding activity of amino-substituted bisphosphonic acids)

RN 63132-38-7 CAPLUS

CN Phosphonic acid, P,P'-[1-amino-3-(dimethylamino)propylidene]bis-(CA INDEX NAME)

PO3H2 H₂N-C-CH₂-CH₂-NMe₂ PO3H2

ACCESSION NUMBER:

1996:642672 CAPLUS Full-text

DOCUMENT NUMBER:

125:316217

TITLE:

Dissociation of binding and antiresorptive properties

of hydroxybisphosphonates by substitution of the

hydroxyl with an amino group

AUTHOR(S):

Van Beek, Ermond; Lowik, Clemens; Que, Ivo;

Papapoulos, Socrates

CORPORATE SOURCE:

Department Endocrinology and Metabolic Diseases,

University Hospital, Leiden, Neth.

SOURCE:

Journal of Bone and Mineral Research (1996), 11(10),

1492-1497

CODEN: JBMREJ; ISSN: 0884-0431

PUBLISHER: DOCUMENT TYPE: Blackwell Journal

LANGUAGE: English

The purpose of this study was to examine the role of the R1 moiety of bisphosphonates in binding to bone mineral and for antiresorptive action. For this, the R1 chain of three clin. useful hydroxybisphosphonates (etidronate, pamidronate, and olpadronate) was substituted with an amino group. The effects of the amino-substituted bisphosphonates were compared with those of their hydroxy counterparts in a crystal growth assay and in fetal mouse long bone cultures which are representative of bisphosphonate actions in vivo. was found that all three amino-substituted compds. and their hydroxy analogs bound with similar affinity to bone mineral and inhibited the growth of calcium oxalate crystals to the same extent. Surprisingly, the antiresorptive effect of olpadronate was totally abolished by the amino substitution of the hydroxyl group while that of pamidronate was reduced by about six-fold and that of etidronate did not change. These studies demonstrate the involvement of the entire bisphosphonate mol. in the cellular mechanism of antiresorptive action. In addition, the amino-substituted analog of olpadronate, which lacks any antiresorptive action but retains all other properties of olpadronate, provides an excellent tool for the study of specific cellular effects involved in bisphosphonate action.

63132-38-7 IT

> RL: BAC (Biological activity or effector, except adverse); BPR (Biological process); BSU (Biological study, unclassified); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses) (dissociation of bone mineral binding and antiresorptive properties of

hydroxybisphosphonates by substitution of hydroxyl with amino group)

RN63132-38-7 CAPLUS

CN Phosphonic acid, P, P'-[1-amino-3-(dimethylamino)propylidene]bis- (CA INDEX NAME)

L38 ANSWER 11 OF 11 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation on

ACCESSION NUMBER:

1999:432676 BIOSIS Full-text

DOCUMENT NUMBER:

PREV199900432676

TITLE:

Differential effects of olpadronate and its

aminosubstituted analog IG-9402 on the regulation of

cytosolic calcium in cultured rat osteoblasts.

AUTHOR(S):

Vazquez, G. [Reprint author]; Boland, R.

[Reprint author]; Roldan, E.; Perez-Lloret,

Α.

CORPORATE SOURCE:

Dept. Biologia, Bioquimica and Farmacia, Universidad

Nacional del Sur, Bahia Blanca, Argentina

SOURCE:

Journal of Bone and Mineral Research, (Sept., 1999) Vol.

14, No. SUPPL. 1, pp. S239. print.

Meeting Info.: Twenty-First Annual Meeting of the American Society for Bone and Mineral Research. St. Louis, Missouri, USA. September 30-October 4, 1999. American Society for

Bone and Mineral Research.

CODEN: JBMREJ. ISSN: 0884-0431.

DOCUMENT TYPE:

Conference; (Meeting)

Conference; Abstract; (Meeting Abstract)

LANGUAGE:

English

ENTRY DATE:

Entered STN: 18 Oct 1999

Last Updated on STN: 3 May 2000

=> fil reg; d stat que 113

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```
L11 1 SEA FILE=REGISTRY ABB=ON "PHOSPHONIC ACID, P,P'-(1-AMINO-3-(DI METHYLAMINO) PROPYLIDENE) BIS-"/CN
L12 2 SEA FILE=REGISTRY ABB=ON 63132-38-7/CRN
L13 3 SEA FILE=REGISTRY ABB=ON (L11 OR L12)
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=> d ide 113 1-3

L13 ANSWER 1 OF 3 REGISTRY COPYRIGHT 2007 ACS on STN

RN 586348-26-7 REGISTRY

ED Entered STN: 16 Sep 2003

CN Phosphonic acid, [1-amino-3-(dimethylamino)propylidene]bis-, nitrate (9CI) (CA INDEX NAME)

MF C5 H16 N2 O6 P2 . x H N O3

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER

CM 1

CRN 63132-38-7 CMF C5 H16 N2 O6 P2

CM

CRN 7697-37-2 CMF H N O3

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L13 ANSWER 2 OF 3 REGISTRY COPYRIGHT 2007 ACS on STN

RN 373645-02-4 REGISTRY

EDEntered STN: 05 Dec 2001

CNPhosphonic acid, [1-amino-3-(dimethylamino)propylidene]bis-, tetrasodium salt (9CI) (CA INDEX NAME)

MF C5 H16 N2 O6 P2 . 4 Na

SR

LCSTN Files: CA, CAPLUS, TOXCENTER, USPAT2, USPATFULL

CRN (63132-38-7)

.
$$PO_3H_2$$

 $H_2N = C + CH_2 + CH_2 + NMe_2$
 PO_3H_2

4 Na

- 1 REFERENCES IN FILE CA (1907 TO DATE)
- 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L13 ANSWER 3 OF 3 REGISTRY COPYRIGHT 2007 ACS on STN

RN 63132-38-7 REGISTRY

ED Entered STN: 16 Nov 1984

Phosphonic acid, P,P'-[1-amino-3-(dimethylamino)propylidene]bis-(CA INDEX NAME)

OTHER CA INDEX NAMES:

Phosphonic acid, [1-amino-3-(dimethylamino)propylidene]bis- (9CI) OTHER NAMES:

CNIG 9402

Lidadronic acid CN

MF C5 H16 N2 O6 P2

CI COM

LCSTN Files: BEILSTEIN*, BIOSIS, CA, CAPLUS, CASREACT, DDFU, DRUGU, IFICDB, IFIPAT, IFIUDB, PHAR, TOXCENTER, USAN, USPAT2, USPATFULL (*File contains numerically searchable property data)

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- 22 REFERENCES IN FILE CA (1907 TO DATE)
- 3 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
- 22 REFERENCES IN FILE CAPLUS (1907 TO DATE)

STRUCTURE SEARCH

=> fil capl; d que 122; s 122 not 123; fil biosis; d que 129; s 129 not 130; fil phar; d que 121
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| L11 | 1 | SEA FILE=REGISTRY | ABB=ON "PHOSPHONIC ACID, | P, P'-(1-AMINO-3-(DI |
|-----|----|---------------------|--------------------------|----------------------|
| | • | METHYLAMINO) PROPYI | LIDENE) BIS-"/CN | |
| L12 | 2 | SEA FILE=REGISTRY | ABB=ON 63132-38-7/CRN | |
| L13 | 3 | SEA FILE=REGISTRY | ABB=ON (L11 OR L12) | • |
| L22 | 24 | SEA FILE=CAPLUS A | BB=ON L13 | |

L39 14 L22 NOT L23

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FILE COVERS 1926 TO DATE.
CAS REGISTRY NUMBERS AND CHEMICAL NAMES (CNs) PRESENT FROM JANUARY 1926 TO DATE.

RECORDS LAST ADDED: 29 August 2007 (20070829/ED)

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| L11 | 1 SEA FILE=REGISTRY ABB=ON "PHOSPHONIC ACID, P,P'-(1-AMINO-3-(DI |
|-----|--|
| | METHYLAMINO) PROPYLIDENE) BIS-"/CN |
| L12 | 2 SEA FILE=REGISTRY ABB=ON 63132-38-7/CRN |
| L13 | 3 SEA FILE=REGISTRY ABB=ON (L11 OR L12) |
| L29 | 2 SEA FILE=BIOSIS ABB=ON L13 |

L40 0 L29 NOT L30

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FILE RELOADED January 2007
FILE LAST UPDATED: Sep 3, 2007 (20070903/ED)

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L20 1 SEA FILE=REGISTRY ABB=ON 63132-38-7/RN

L21 1 SEA FILE=PHAR ABB=ON L20

=> dup rem 139,121

DUPLICATE IS NOT AVAILABLE IN 'PHAR'.

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15 DUP REM L39 L21 (0 DUPLICATES REMOVED) ANSWERS '1-14' FROM FILE CAPLUS

ANSWER '15' FROM FILE PHAR

=> d ibib abs hitind 1-14; d all 15; fil hom

L41 ANSWER 1 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2006:818726 CAPLUS Full-text

DOCUMENT NUMBER: 146:288334

TITLE: Dissociation of the pro-apoptotic effects of

bisphosphonates on osteoclasts from their

anti-apoptotic effects on osteoblasts/osteocytes with

novel analogs

AUTHOR(S): Plotkin, Lilian I.; Manolagas, Stavros C.; Bellido,

Teresita

CORPORATE SOURCE:

Division of Endocrinology and Metabolism, The Center for Osteoporosis and Metabolic Bone Diseases, The

Central Arkansas Veterans Healthcare System,

University of Arkansas for Medical Sciences, Little

Rock, AR, 72205, USA

SOURCE:

Bone (San Diego, CA, United States) (2006), 39(3),

443-452

CODEN: BONEDL; ISSN: 8756-3282

PUBLISHER: DOCUMENT TYPE:

Elsevier Journal

DOCUMENT TYPE: Journal LANGUAGE: English

AB Bisphosphonates induce osteoclast apoptosis, thereby decreasing bone resorption and reducing the rate of bone remodeling. Earlier work from our group and others has demonstrated that, addnl., bisphosphonates prevent osteoblast and osteocyte apoptosis in vivo and in vitro, raising the possibility that perhaps part of their anti-fracture efficacy may result from preserving the integrity of the osteocyte network and prolonging the working time of bone forming cells. Whereas induction of osteoclast apoptosis results from inhibition of the mevalonate pathway or from conversion to toxic ATP analogs, prevention of osteoblastic cell apoptosis is mediated by connexin43 hemichannel opening and activation of the extracellular signal-regulated kinases (ERKs). We examined here the ability of several bisphosphonates, including novel analogs, to exert these two effects. All 16 bisphosphonates studied inhibited etoposide-induced apoptosis of MLO-Y4 osteocytic cells and osteoblastic cells derived from calvaria, with EC50 between 10-12 and 10-10 M. On the other hand, only 10 analogs induced apoptosis of RAW-264.7-cell-derived osteoclasts. Each of the 6 bisphosphonates that lack pro-apoptotic activity in osteoclasts but retain anti-apoptotic activity in osteoblasts and osteocytes has a structural-related analog that is active in both cell types. These findings indicate that the structural prerequisites for the antiapoptotic effect of bisphosphonates on cells of the osteoblastic lineage are less stringent than the ones required to induce osteoclast apoptosis and confirm that bisphosphonates act on the two cell types by distinct mechanisms. Preservation of osteoblast and osteocyte viability without inducing osteoclast apoptosis by these bisphosphonates analogs opens new possibilities for the treatment of bone fragility in conditions in which a decrease in bone remodeling is not desirable.

CC 1-12 (Pharmacology)

IT 63132-38-7, IG9402

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(bisphosphonate analog IG9402 prevented osteoblast and osteocyte apoptosis without affecting mouse osteoclasts)

REFERENCE COUNT:

48 THERE ARE 48 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L41 ANSWER 2 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2003:986477 CAPLUS Full-text

DOCUMENT NUMBER:

140:156750

TITLE:

Quantitative Structure-Activity Relationships for

 $\gamma\delta$ T Cell Activation by Bisphosphonates

AUTHOR (S):

Sanders, John M.; Ghosh, Subhash; Chan, Julian M. W.; Meints, Gary; Wang, Hong; Raker, Amy M.; Song,

Yongcheng; Colantino, Alison; Burzynska, Agnieszka; Kafarski, Pawel; Morita, Craig T.; Oldfield, Eric Department of Chemistry, University of Illinois at

CORPORATE SOURCE:

Urbana-Champaign, Urbana, IL, 61801, USA

SOURCE:

Journal of Medicinal Chemistry (2004), 47(2), 375-384

CODEN: JMCMAR; ISSN: 0022-2623

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 140:156750

γδ T cells are the first line of defense against many infectious organisms and AB are also involved in tumor cell surveillance and killing. They are stimulated by a broad range of small, phosphorus-containing antigens (phosphoantigens) as well as by the bisphosphonates commonly used in bone resorption therapy, such as pamidronate and risedronate. Here, we report the activation of $\gamma\delta$ T cells by a broad range of bisphosphonates and develop a pharmacophore model for $\gamma\delta$ T cell activation, in addition to using a comparative mol. similarity index anal. (CoMSIA) approach to make quant. relationships between $\gamma\delta$ T cell activation by bisphosphonates and their three-dimensional structures. CoMSIA analyses yielded R2 values of .apprx.0.8-0.9 and q2 values of .apprx.0.5-0.6 for a training set of 45 compds. Using an external test set, the activities (IC50 values) of 16 compds. were predicted within a factor of 4.5, on average The CoMSIA fields consisted of .apprx.40% hydrophobic, .apprx.40% electrostatic, and .apprx.20% steric interactions. Since bisphosphonates are known to be potent, nanomolar inhibitors of the mevalonate/isoprene pathway enzyme farnesyl pyrophosphate synthase (FPPS), we also compared the pharmacophores for $\gamma\delta$ T cell activation with those for FPPS inhibition, using the Catalyst program. The pharmacophores for $\gamma\delta$ T cell activation and FPPS inhibition both consisted of two neg. ionizable groups, a pos. charge feature and an endocyclic carbon feature, all having very similar spatial dispositions. In addition, the CoMSIA fields were quite similar to those found for FPPS inhibition by bisphosphonates. The activities of the bisphosphonates in $\gamma\delta$ T cell activation were highly correlated with their activities in FPPS inhibition: R = 0.88, p = 0.002, vs. a human recombinant FPPS (N = 9 compds.); R = 0.82, p < 0.0001, for an expressed Leishmania major FPPS (N = 45 compds.). The bisphosphonate $\gamma\delta$ T cell activation pharmacophore differs considerably, however, from that reported previously for γδ T cell activation by phosphoantigens (Gossman, W.; Oldfield, E. J. Med. Chemical 2002, 45, 4868-4874), suggesting different primary targets for the two classes of compds. The ability to quite accurately predict the activity of bisphosphonates as γδ T cell activators by using 3D QSAR techniques can be expected to help facilitate the design of addnl. bisphosphonates for potential use in immunotherapy.

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CC 1-3 (Pharmacology)
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IT
     2809-22-5
                 16559-82-3
                              32545-64-5
                                           32545-65-6
                                                         32545-72-5
                                                                      32579-17-2
     40391-99-9
                  56152-34-2
                              56375-74-7 63132-38-7
                                                        67242-32-4
                                            70010-79-6
     70010-75-2
                  70010-76-3
                               70010-77-4
                                                          70010-82-1
     70010-83-2
                  70010-87-6
                               71066-25-6
                                            71066-28-9
                                                          71066-29-0
     71066-40-5
                  79778-41-9
                               89732-96-7
                                            104261-69-0
                                                           105462-22-4
     105462-23-5
                   105462-24-6
                                 105462-25-7
                                                111072-49-2
                                                              114084-78-5
     114119-81-2
                   118072-93-8, Zometa
                                          121368-58-9, Olpadronate
                                                                     124351-85-5
     124351-86-6
                   124351-87-7
                                 124369-71-7
                                                129318-43-0, Fosamax
     129951-00-4
                   134579-56-9
                                 149543-15-7
                                                180064-38-4
                                                              634612-13-8
     634612-14-9
                   634612-16-1
                                 656238-26-5
    RL: DMA (Drug mechanism of action); PAC (Pharmacological activity); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
```

(quant. structure-activity relationships for $\gamma\delta$ T cell activation by bisphosphonates)

REFERENCE COUNT: 49 THERE ARE 49 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L41 ANSWER 3 OF 15 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 2003:652131 CAPLUS Full-text DOCUMENT NUMBER: 139:214237

TITLE:

Preparation of nitrate prodrugs able to release nitric oxide in a controlled and selective way and their use for prevention and treatment of inflammatory, ischemic

and proliferative diseases

INVENTOR (S):

Scaramuzzino, Giovanni

PATENT ASSIGNEE(S):

Italy

SOURCE:

Eur. Pat. Appl., 313 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE -----------EP 1336602 A1 20030820 EP 2002-425075

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,

IE, SI, LT, LV, FI, RO, MK, CY, AL, TR

PRIORITY APPLN. INFO.:

EP 2002-425075

20020213

GI

AΒ New pharmaceutical compds. of general formula F-(X)q (I) [q = 1-5, preferably]1; F is chosen among drugs such as δ -tocopherol, clidanac, diethylhomospermine, glucosamine, thymocartin, vofopitant, etc.; X is chosen among 4 groups M, T, V, and Y where M = ONO2, nitrate salt, nitrite ester, ONO, thoinitrite, SNO, etc., T = OR1-M, OR1OR1-M, SR1NR2R1-M, NR2R1-M, NR2R1SR1-M, etc., R1 = saturated or unsatd., linear or branched alkylene, having 1 to 21 carbon atoms or a saturated or unsatd., optionally heterosubstituted or branched cycloalkylene, having 3 to 7 carbon atoms or an optionally heterosubstituted arylene having 3 to 7 carbon atoms; R2 = H, saturated or unsatd., linear or branched 1-21 carbon atom alkyl, saturated or unsatd. optionally heterosubstituted or branched 3-7 carbon cycloalkyl, optionally heterosubstituted 3-7 carbon aryl; R1, R2 = OH, SH, F, Cl, Br, OPO3H2, CO2H, etc.; bond between F and T = carboxylic ester, carboxylic amide, glycoside, azo, thioester, sulfonic ester, etc.; V = Z-M2, OZ-M2, NR2Z-M2, R1Z-M2, OR1-M2, OR1Z-M2, M2 = M, R1-M, OR1-M, SR1-M, NR2R1-M; ZM2 =COCH2CH(M2)CH2N+Me3, COCH2CH2COM2, COCH(NHR2)CH2M2, etc.; Y = 4-COC6H4CH2ONO2, O(CH2)40NO2, COCH(NH2)CH20NO2, 3-OC6H4CH20NO2, etc.] were prepared For example, α -tocopherol reacted with 4-HO2CC6H4CH2ONO2 to give the nitroxymethyl derivative II. The compds. of general formula I are nitrate prodrugs which can release nitric oxide in vivo in a controlled and selective way and without hypotensive side effects and for this reason they are useful for the preparation of medicines for prevention and treatment of inflammatory, ischemic, degenerative and proliferative diseases of musculoskeletal, tegumental, respiratory, gastrointestinal, genito-urinary and central nervous systems.

ICM C07C205-00

IC

ICS A61K031-00

CC 26-1 (Biomolecules and Their Synthetic Analogs) Section cross-reference(s): 1, 28, 29, 33, 34, 63 IT 13005-09-9P 96513-33-6P 116539-59-4P 198483-54-4P 257625-98-2P 329976-33-2P 352464-98-3P 398454-56-3P 398460-42-9P 410071-16-8P 571186-52-2P 586347-21-9P 586347-23-1P 586347-25-3P 586347-26-4P 586347-28-6P 586347-29-7P 586347-31-1P 586347-32-2P 586347-33-3P 586347-34-4P 586347-36-6P 586347-38-8P 586347-39-9P 586347-42-4P 586347-43-5P 586347-45-7P 586347-46-8P 586347-47-9P 586347-48-0P 586347-50-4P 586347-51-5P 586347-52-6P 586347-53-7P 586347-54-8P 586347-55-9P 586347-56-0P 586347-57-1P 586347-60-6P 586347-58-2P 586347-62-8P 586347-63-9P 586347-64-0P 586347-65-1P 586347-66-2P 586347-68-4P 586347-69-5P 586347-70-8P 586347-72-0P 586347-73-1P 586347-75-3P 586347-76-4P 586347-77-5P 586347-79-7P 586347-80-0P 586347-81-1P 586347-82-2P 586347-86-6P 586347-92-4P 586347-94-6P 586347-95-7P 586347-97-9P 586348-00-7P 586348-01-8P 586348-03-0P 586348-04-1P 586348-05-2P 586348-07-4P 586348-08-5P 586348-09-6P 586348-10-9P 586348-11-0P 586348-12-1P 586348-13-2P 586348-14-3P 586348-15-4P 586348-16-5P 586348-17-6P 586348-18-7P 586348-19-8P 586348-20-1P 586348-22-3P 586348-23-4P 586348-24-5P 586348-25-6P 586348-26-7P 586348-27-8P 586348-28-9P 586348-29-0P 586348-30-3P 586348-31-4P 586348-32-5P 586348-34-7P 586348-35-8P 586348-36-9P 586348-37-0P 586348-39-2P 586348-40-5P 586348-42-7P 586348-43-8P 586348-45-0P 586348-47-2P 586348-49-4P 586348-51-8P 586348-52-9P 586348-53-0P 586348-55-2P 586348-57-4P 586348-59-6P 586348-61-0P 586348-62-1P 586348-63-2P 586348-64-3P 586348-65-4P 586348-66-5P 586348-67-6P 586348-68-7P 586348-69-8P 586348-70-1P 586348-71-2P 586348-72-3P 586348-73-4P 586348-74-5P 586348-75-6P 586348-76-7P 586348-77-8P 586348-78-9P 586348-79-0P 586348-80-3P 586348-82-5P 586348-84-7P 586348-85-8P 586348-86-9P 586348-87-0P 586348-88-1P 586348-90-5P 586348-91-6P 586348-92-7P 586348-93-8P 586348-95-0P 586348-97-2P 586348-98-3P 586348-99-4P 586349-00-0P 586349-02-2P 586349-04-4P 586349-05-5P 586349-06-6P 586349-08-8P 586349-10-2P 586349-09-9P 586349-12-4P 586349-13-5P 586349-14-6P 586349-15-7P 586349-16-8P 586349-17-9P 586349-18-0P 586349-19-1P 586349-21-5P 586349-20-4P 586349-23-7P 586349-22-6P 586349-24-8P 586349-25-9P 586349-26-0P 586349-27-1P 586349-28-2P 586349-29-3P 586349-30-6P 586349-31-7P 586349-32-8P 586349-33-9P 586349-34-0P 586349-36-2P 586349-37-3P 586349-38-4P 586349-39-5P 586349-40-8P 586349-41-9P 586349-43-1P 586349-44-2P 586349-45-3P 586349-46-4P 586349-47-5P 586349-48-6P 586349-49-7P 586349-50-0P 586349-51-1P 586349-52-2P 586349-53-3P 586349-54-4P 586349-55-5P 586349-56-6P 586349-57-7P 586349-58-8P 586349-60-2P 586349-61-3P 586349-63-5P 586349-64-6P 586349-66-8P 586349-67-9P 586349-69-1P 586349-70-4P 586349-71-5P 586349-72-6P 586349-74-8P 586349-76-0P 586349-77-1P 586349-79-3P 586349-80-6P 586349-81-7P 586349-83-9P 586349-84-0P 586349-85-1P 586349-86-2P 586349-87-3P 586349-88-4P 586349-89-5P 586349-90-8P 586349-91-9P 586349-92-0P 586349-93-1P 586349-94-2P 586349-95-3P 586349-96-4P 586349-97-5P 586349-98-6P 586349-99-7P 586350-01-8P 586350-02-9P 586350-03-0P 586350-04-1P 586350-05-2P RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(preparation of nitrate prodrugs for treating or preventing inflammatory, ischemic, degenerative, and proliferative diseases)

REFERENCE COUNT: 19 THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L41 ANSWER 4 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:539062 CAPLUS Full-text

DOCUMENT NUMBER:

137:226194

TITLE:

Highly Potent Geminal Bisphosphonates. From

Pamidronate Disodium (Aredia) to Zoledronic Acid

AUTHOR (S):

Widler, Leo; Jaeggi, Knut A.; Glatt, Markus; Mueller, Klaus; Bachmann, Rolf; Bisping, Michael; Born, Anne-Ruth; Cortesi, Reto; Guiglia, Gabriela; Jeker, Heidi; Klein, Remy; Ramseier, Ueli; Schmid, Johann; Schreiber, Gerard; Seltenmeyer, Yves; Green, Jonathan

CORPORATE SOURCE:

Arthritis and Bone Metabolism Therapeutic Area, Novartis Pharma Research, Basel, CH-4002, Switz.

SOURCE:

Journal of Medicinal Chemistry (2002), 45(17),

3721-3738

CODEN: JMCMAR; ISSN: 0022-2623

PUBLISHER:

American Chemical Society

DOCUMENT TYPE:

Journal English

LANGUAGE: OTHER SOURCE(S):

CASREACT 137:226194

AB

Bisphosphonates (BPs) are pyrophosphate analogs in which the oxygen in P-O-P has been replaced by a carbon, resulting in a metabolically stable P-C-P structure. Pamidronate (1b, Novartis), a second-generation BP, was the starting point for extensive SAR studies. Small changes of the structure of pamidronate lead to marked improvements of the inhibition of osteoclastic resorption potency. Alendronate (1c, MSD), with an extra methylene group in the N-alkyl chain, and olpadronate (1h, Gador), the N,N-di-Me analog, are about 10 times more potent than pamidronate. Extending one of the N-Me groups of olpadronate to a pentyl substituent leads to ibandronate (1k, Roche, Boehringer-Mannheim), which is the most potent close analog of pamidronate. Even slightly better antiresorptive potency is achieved with derivs. having a Ph group linked via a short aliphatic tether of three to four atoms to nitrogen, the second substituent being preferentially a Me group (e.g., 4g, 4j, 5d, or 5r). The most potent BPs are found in the series containing a heteroarom. moiety (with at least one nitrogen atom), which is linked via a single methylene group to the geminal bisphosphonate unit. Zoledronic acid (6i), the most potent derivative, has an ED50 of 0.07 mg/kg in the TPTX in vivo assay after s.c. administration. It not only shows by far the highest therapeutic ratio when comparing resorption inhibition with undesired inhibition of bone mineralization but also exhibits superior renal tolerability. Zoledronic acid (6i) has thus been selected for clin. development under the registered trade name Zometa. The results of the clin. trials indicate that low doses are both efficacious and safe for the treatment of tumor-induced hypercalcemia, Paget's disease of bone, osteolytic metastases, and postmenopausal osteoporosis.

CC 1-3 (Pharmacology)

TΤ 29712-30-9P 32545-72-5P 56152-35-3P 63132-38-7P 63161-30-8P 66376-36-1P, Alendronate 63132-40-1P 67242-32-4P 79778-41-9P, Neridronate 86235-67-8P 89732-96-7P 104261-68-9P 114084-78-5P, Ibandronate 114084-82-1P 114119-81-2P 116162-22-2P 116786-78-8P 116786-79-9P 116786-83-5P 116786-85-7P 116786-88-0P 116786-89-1P 116786-90-4P 118054-12-9P 118054-15-2P 118054-16-3P 118054-18-5P 118054-23-2P 118054-19-6P 118054-20-9P 118054-31-2P 118054-32-3P 118054-33-4P 118054-41-4P 118054-42-5P 118054-51-6P 118054-52-7P 118072-93-8P 118694-16-9P 121368-58-9P, Olpadronate 124351-85-5P 124369-71-7P 124369-72-8P 124369-73-9P 124369-77-3P 124369-80-8P 124369-81-9P 124369-83-1P 125946-91-0P 128202-57-3P 129951-00-4P 129951-01-5P 129951-02-6P 131654-39-2P 131654-40-5P 131654-41-6P 131654-58-5P 132423-84-8P 132423-86-0P 132423-87-1P 132423-88-2P 132423-89-3P 132423-90-6P 132423-92-8P 132423-94-0P 132423-95-1P 132423-96-2P 132423-97-3P 132423-98-4P 132423-99-5P

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132424-00-1P
               132424-01-2P
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                                             134579-55-8P
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136671-90-4P
               142830-99-7P
                              149226-80-2P
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183446-98-2P
               209002-31-3P
                              209002-32-4P
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459870-47-4P
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                                             459870-50-9P
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459870-52-1P
               459870-53-2P
                              459870-54-3P
                                             459870-55-4P
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               459870-58-7P
459870-57-6P
                              459870-59-8P
                                             459870-60-1P
                                                            459870-61-2P
RL: ADV (Adverse effect, including toxicity); PAC (Pharmacological
activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL
(Biological study); PREP (Preparation); USES (Uses)
   (bisphosphonates preparation and structure-related bone antiresorptive
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properties) REFERENCE COUNT: THERE ARE 88 CITED REFERENCES AVAILABLE FOR THIS 88

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L41 ANSWER 5 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER:

2001:833023 CAPLUS Full-text

DOCUMENT NUMBER:

135:376738

TITLE:

Compounds and methods for modulating cerebral amyloid

angiopathy using inhibitors of an amyloid β

peptide

INVENTOR(S):

Green, Allan M.; Gervais, Francine

PATENT ASSIGNEE(S): SOURCE:

Neurochem, Inc., Can.

PCT Int. Appl., 68 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| | PATENT NO. | | | | | | | | | DATE | | | | | | | | |
|-----------|------------------------|------|------|-----|-----|----------|------|------|-----|----------------|------|------|-----|-----|----------|------|-----|--|
| | 2001 | | | | | | | | | WO 2000-IB2078 | | | | | 20001222 | | | |
| WO | 2001 | 0850 | 93 | | A3 | 20020829 | | | | | | | | | | | | |
| WO | 2001 | 0850 | 93 . | | Α9 | | 2002 | 0926 | | | • | | | | | | | |
| | W: | ΑE, | AG, | AL, | AM, | AT, | AU, | AZ, | BA, | BB, | ВG, | BR, | BY, | ΒZ, | CA, | CH, | CN, | |
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| | | LU, | LV, | MA, | MD, | MG, | MK, | MN, | MW, | MX, | MZ, | NO, | NZ, | PL, | PT, | RO, | RU, | |
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| : | RW: | GH; | GM, | ΚE, | LS, | MW, | MZ, | SD, | SL, | SZ, | TZ, | UG, | ZW, | AT, | BE, | CH, | CY, | |
| | | | | | | | GB, | | | | | | | | | | | |
| | | | | | | | GA, | | | | | | | | | • | • | |
| CA | 2395 | | | | | | | | | | | | | | | 0001 | 222 | |
| AU | 2001 | 8431 | 3 | | Α | | 2001 | 1120 | | AU 2 | 001- | 8431 | 3 | | 2 | 0001 | 222 | |
| | 1251 | | | | | | | | | | | | | | | 0001 | | |
| | | | | | | | ES, | | | | | | | | | MC, | PT. | |
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| BR | 2000 | 0166 | 52 | | Α | | 2002 | 1119 | | BR 2 | -000 | 1665 | 2 | | 2 | 0001 | 222 | |
| | 2003 | | | | | | 2003 | | | | | | | | | 0001 | | |
| US | 6670 | 399 | | | B2 | | 2003 | 1230 | | | | | | | | | | |
| JP | 2003 | | | | | | 2003 | 1105 | 1 | JP 2 | 001- | 5817 | 48 | | 2 | 0001 | 222 | |
| | 2002 | | | | | | | | | | | | | | | 0020 | 621 | |
| AU | 2006 | 2014 | 45 | | A1 | | 2006 | 0504 | | AU 2 | 006- | 2014 | 45 | | 2 | 0060 | 406 | |
| PRIORIT | PRIORITY APPLN. INFO.: | | | . : | | | | | | US 1 | 999- | 1718 | 77P | 3 | P 1 | 9991 | 223 | |
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OTHER SOURCE(S): MARPAT 135:376738

The invention provides methods of inhibiting cerebral amyloid angiopathy (CAA) and treating a disease state characterized by cerebral amyloid angiopathy,

e.g., Alzheimer's disease, in a subject using an inhibitor of the 39-40 amino acid amyloid β peptide (A β 40). The A β 40 inhibitor is selected from, e.g., sulfonic acid derivs., such as ethanesulfonic acid, 1,2-ethanedisulfonic acid, 1-propanesulfonic acid, 1,3-propanedisulfonic acid, 1,4-butanedisulfonic acid, 1,5-pentanedisulfonic acid, 2-aminoethanesulfonic acid, 4-hydroxy-1butanesulfonic acid, 1-butanesulfonic acid, 1-decanesulfonic acid, 2propanesulfonic acid, 3-pentanesulfonic acid, 4-heptanesulfonic acid, etc., and pharmaceutically acceptable salts thereof or from from phosphonic acid derivs., such as diethylphosphonoacetic acid, phenylphosphonic acid, 3aminopropylphosphonic acid, propylphosphonic acid, etc. The compds. are formulated in a dispersion system, a liposome formulation, or microspheres using a polymeric matrix. The polymeric matrix is selected from natural polymers, such as albumin, alginate, cellulose derivs., collagen, fibrin, gelatin, and polysaccharides, or synthetic polymers such as polyesters, polyethylene glycol, poloxamers, and polyanhydrides. For example, the ability of compds. of the invention to inhibit CAA was measured in 9 wk old hAPP transgenic mice treated with two different concns. of a compound of the present invention, 3-amino-1-propanesulfonic acid sodium salt, 100 and 30 mg/kg. Mice were administered the compound for 8 wk, after which they were sacrificed and their brains were perfused and processed for histol, staining with Thioflavin S. This method may also be used as a screening method for determining activity of a candidate compound for inhibiting CAA. The extent of CAA in brain sections obtained from these animals was qual. determined following staining. The results indicate that the test compound was effective in (i) reducing the number of mice showing CAA, and (ii) showing an effect on the severity of the deposition seen in the brain vasculature of these animals.

IC ICM A61K

CC 63-6 (Pharmaceuticals)

Section cross-reference(s): 1

IT107-35-7, 2-Aminoethanesulfonic acid 110-04-3, 1,2-Ethanedisulfonic acid 116-63-2 149-45-1 288-94-8, 1H-Tetrazole 594-45-6, Ethanesulfonic acid 831-59-4 860-22-0 926-39-6 Methylphosphonic acid 1068-21-9, Diethyl phosphoramidate 1071-83-6, N-Phosphonomethylglycine 1120-71-4 1132-61-2, 4-Morpholinepropanesulfonic acid 1135-40-6 1571-33-1, Phenylphosphonic 1633-83-6 2386-47-2, 1-Butanesulfonic acid 3687-18-1, 3-Amino-1-3095-95-2, Diethylphosphonoacetic acid propanesulfonic acid 4408-78-0, Phosphonoacetic acid 4672-38-2, Propylphosphonic acid 4923-84-6 5117-07-7 5284-66-2. 1-Propanesulfonic acid 5399-58-6 5652-28-8 5994-73-0 6779-09-5, Ethylphosphonic acid 7365-45-9 13138-33-5, 3-Aminopropylphosphonic 13991-98-5, 14047-23-5; (1-Aminopropyl) phosphonic acid 13419-61-9 14159-48-9, 2-Propanesulfonic acid acid 14650-46-5 15471-17-7 15763-57-2 18039-42-4 20283-21-0, 1-Decanesulfonic acid 21668-77-9, 1,3-Propanedisulfonic acid 23052-80-4 23052-81-5 25331-57-1 25595-59-9 26978-64-3, 4-Hydroxy-1-butanesulfonic acid 27665-39-0, 1,4-Butanedisulfonic acid 27797-35-9 31465-25-5 34159-44-9 36585-99-6 37810-68-7 38911-09-0 40391-99-9 40465-65-4. N-Phosphonomethylglycine trisodium salt 51224-03-4 51224-04-5 51650-30-7, 3-Pentanesulfonic acid 51762-95-9 53329-36-5 63585-09-1, Phosphonoformic acid trisodium salt 58849-79-9 60142-96-3 75277-39-3 71119-22-7 72217-85-7 73858-58-9 76326-31-3, 2-Amino-5-phosphonopentanoic acid 78739-01-2, D-(-)-2-Amino-4phosphonobutanoic acid 79055-67-7 79055-68-8 81338-23-0 81338-24-1, L-(+)-2-Amino-7-phosphonoheptanoic acid 82283-67-8 82283-68-9 82977-27-3 87625-44-3 88246-85-9 91357-22-1 91586-81-1 99107-93-4 101020-77-3, 1,5-Pentanedisulfonic acid 102805-84-5 108084-41-9 112980-83-3 117414-74-1 126453-07-4 128241-72-5 129318-43-0 131177-53-2 138199-51-6 143018-67-1 145544-51-0 157381-42-5 168977-94-4,

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3-Aminopropyl (methyl) phosphinic acid hydrochloride
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4-Heptanesulfonic acid
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                             373645-10-4
                                            373645-11-5
                                                          373645-12-6
373645-13-7
```

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(inhibitors of amyloid β peptide for modulating cerebral amyloid angiopathy)

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L41 ANSWER 6 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:416728 CAPLUS Full-text
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DOCUMENT NUMBER: 135:14356

TITLE: Phosphonate compounds, and preparation thereof, for

treating medical disorders

INVENTOR(S): Hostetler, Karl Y.; Beadle, James R.; Kini, Ganesh D.

PATENT ASSIGNEE(S): The Regents of the University of California, San

Diego, USA

SOURCE: PCT Int. Appl., 47 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE: En

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO | KIND | | DATE | | i | APPLICATION NO. | | | | | DATE | | | | | |
|-----------|--------|-----|------------|-----|------|-----------------|-----|------|-------|------|------|-----|-----|-------|-----|--|
| | | | - | | | | | | | | | | | | | |
| WO 200103 | 9724 | | A2 | : | 2001 | 0607 | 1 | WO 2 | 7-00C | JS33 | 079 | | 20 | 00012 | 204 | |
| WO 200103 | 9724 | | A 3 | | 2001 | 1018 | | | | | | | | | | |
| W: A | E, AG, | AL, | AM, | AT, | AU, | AZ, | BA, | BB, | BG, | BR, | BY, | ΒZ, | CA, | CH, | CN, | |
| · C | R, CU, | CZ, | DE, | DK, | DM, | DZ, | EE, | ES, | FI, | GB, | GD, | GE, | GH, | GM, | HR, | |
| . H | U, ID, | IL, | IN, | IS, | JP, | KE, | KG, | KP, | KR, | ΚZ, | LC, | LK, | LR, | LS, | LT, | |
| L | U, LV, | MA, | MD, | MG, | MK, | MN, | MW, | MX, | MZ, | NO, | NZ, | PL, | PT, | RO, | RU, | |
| S | D, SE, | SG, | SI, | SK, | SL, | TJ, | TM, | TR, | TT, | TZ, | UΑ, | UG, | US, | UZ, | VN, | |
| Y | U, ZA, | zw | | | | | | | | | | | | | | |

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RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
     CA 2393410
                          A1
                                20010607 CA 2000-2393410
    AU 200119497
                                            AU 2001-19497
                          Α
                                20010612
                                                                   20001204
    AU 785355
                          B2
                                20070201
    EP 1233770
                          A2
                                20020828
                                            EP 2000-982468
                                                                   20001204
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             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
    BR 2000016058
                         Α
                                20030715
                                            BR 2000-16058
                                                                   20001204
    JP 2004500352
                          T ·
                                20040108
                                            JP 2001-541459
                                                                   20001204
    RU 2258707
                          C2
                                20050820
                                            RU 2002-118327
                                                                   20001204
     IN 2002DN00553
                          Α
                                20040228
                                            IN 2002-DN553
                                                                   20020531
    MX 2002PA05490
                         Α
                                20040910
                                            MX 2002-PA5490
                                                                   20020603
    US 2004019232
                         A1
                                20040129
                                            US 2002-148374
                                                                   20021106
    US 6716825
                          B2
                                20040406
    ZA 2002004194
                         Α
                                20030820
                                            ZA 2002-4194
                                                                   20021204
    US 2004127735
                                            US 2004-759345
                         A1
                                20040701
                                                                   20040115
    US 7034014
                         B2
                                20060425
    US 2005176673
                         A1
                                20050811
                                            US 2005-100882
                                                                   20050406
    US 7094772
                         B2
                                20060822
    US 2005182019
                         A1
                                20050818
                                            US 2005-101259
                                                                   20050406
    US 7098197
                         B2
                                20060829
    US 2006281706
                         A1
                                20061214
                                            US 2006-506292
                                                                   20060817
    AU 2006252074
                          A1
                                20070118
                                            AU 2006-252074
                                                                   20061215
    US 2007161602
                                20070712
                         A1
                                            US 2007-715604
                                                                   20070307
                                                                P 19991203
PRIORITY APPLN. INFO.:
                                            US 1999-168813P
                                            US 2000-205719P
                                                                P 20000519
                                            AU 2001-19497
                                                                T0 20001204
                                            WO 2000-US33079
                                                                W 20001204
                                            US 2002-148374
                                                                A1 20021106
                                            US 2004-759345
                                                                A1 20040115
                                            US 2005-100882
                                                                A1 20050406
                                            US 2006-506292
                                                                A1 20060817
```

OTHER SOURCE(S): MARPAT 135:14356

- AB The invention discloses phosphonate compds., compns. containing them, processes for obtaining them, and their use for treating a variety of medical disorders, e.g. osteoporosis and other disorders of bone metabolism, cancer, and viral infections. Preparation of compds. of the invention, e.g. 1-0-hexadecylpropanediol-3-alendronate, is described.
- IC ICM A61K
- CC 1-12 (Pharmacology)
- IT 147-94-4D, Cytosine arabinoside, derivs. 2809-21-4D, derivs.
 4291-63-8D, 2-Chlorodeoxyadenosine, derivs. 10596-23-3D, derivs.
 13598-36-2D, Phosphonic acid, derivs. 21679-14-1D, Fludarabine, derivs.
 30516-87-1D, Azidothymidine, derivs. 38819-10-2D, derivs. 40391-99-9D, derivs. 63132-38-7D, derivs. 66376-36-1D, Alendronate, derivs.
 89987-06-4D, Tiludronate, derivs. 95058-81-4D, Gemcitabine, derivs.
 105462-24-6D, derivs. 106941-25-7D, Adefovir, derivs. 113852-37-2D, Cidofovir, derivs. 114084-78-5D, Ibandronate, derivs. 121368-58-9D, Olpadronate, derivs. 125946-92-1D, EB-1053, derivs. 127757-45-3D, derivs. 147127-20-6D, Tenofovir, derivs.
 RL: BAC (Biological activity or effector, except adverse); BSU (Biological study); USES

(phosphonate compds., and preparation thereof, for treating medical disorders)

L41 ANSWER 7 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2000:351360 CAPLUS Full-text

DOCUMENT NUMBER:

132:343333

TITLE:

Increasing bone strength with selected bisphosphonates

INVENTOR(S): Manolagas, Stavros C.; Bellido, Teresita

PATENT ASSIGNEE(S):

The Board of Trustees for the University of Arkansas,

USA; Gador S.A.

SOURCE:

PCT Int. Appl., 71 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| P | PATENT NO. | | | | | KIND DATE | | APPLICATION NO. | | | | | DATE | | | | |
|--------|------------|--------|------|------|-----|-----------|------|-----------------|-----|-------|-------|-------|------|-----|------|------|-------|
| - | | | | | - | | | | | | | | | - | | | |
| . W | 10 20 | 000289 | 82 | | A2 | | 2000 | 0525 | 1 | WO 1: | 999-1 | US27 | 528 | | 1 | 9991 | 119 |
| W | 70 20 | 000289 | 82 | | А3 | | 2002 | 0711 | | | | | | | | | |
| | W | : AE, | ΑL, | AM, | AT, | AU, | ΑZ, | BA, | BB, | BG, | BR, | BY, | CA, | CH, | CN, | CR, | CU, |
| | | CZ, | DE, | DK, | DM, | EE, | ES, | FI, | GB, | GD, | GE, | GH, | GM, | HR, | HU, | ID, | L, |
| | | IN, | IS, | JP, | KΕ, | KG, | ΚP, | KR, | KZ, | LC, | LK, | LR, | LS, | LT, | LU, | LV, | MA, |
| | | MD, | MG, | MK, | MN, | MW, | MX, | NO, | NZ, | PL, | PT, | RO, | RU, | SD, | SE, | SG, | SI, |
| | | SK, | SL, | ТJ, | TM, | TR, | TT, | UA, | UG, | US, | UZ, | VN, | YU, | ZA, | ZW, | AM, | ΑZ, |
| | | BY, | KG, | KZ, | MD, | RU, | TJ, | TM | | | | | | | | | |
| | RI | N: GH, | GM, | ΚE, | LS, | MW, | SD, | SL, | SZ, | TZ, | UG, | ZW, | AT, | BE, | CH, | CY, | DE, |
| | | DK, | ES, | FI, | FR, | GB, | GR, | ΙE, | IT, | LU, | MC, | NL, | PT, | SE, | BF, | ВJ, | CF, |
| | | CG, | CI, | ·CM, | GA, | GN, | GW, | ML, | MR, | NE, | SN, | TD, | TG | | | | |
| A | U 200 | 000152 | 57 | | Α | | 2000 | 0605 | 7 | AU 2 | 000-1 | 1525 | 7 | | . 1 | 9991 | 119 |
| U | JS 64 | 16737 | | | B1 | | 2002 | 0709 | 1 | US 1 | 999- | 44384 | 41 | | 1 | 9991 | 119 |
| PRIORI | TY A | PPLN. | INFO | .: | | | | | 1 | US 1 | 998- | 1092 | 37P |] | P 1: | 9981 | 119 |
| | | | | | | | | , | 1 | US 1 | 999-: | 1654 | 80P | 1 | P 1 | 9991 | 115 . |
| | | | | | | | | | 1 | WO 1 | 999-1 | US27 | 528 | 7 | W 1: | 9991 | 119 |

- AΒ The present invention is a method and composition to increase bone strength in a manner that decreases fracture incidence, which may or may not include increasing bone mineral d. ("BMD"). The invention includes administering an effective amount of a bisphosphonate to a host in need thereof to increase bone strength, which inhibits the apoptosis of osteoblasts and osteocytes, without a significant effect on osteoclasts. In one embodiment, the bisphosphonate is not 1-amino-3-(N,N-dimethylamino) - propyliden-1,1bisphosphonic acid or its pharmaceutically acceptable salt. An increase in osteoblast life span can lead to an increase in bone mass, i.e., an anabolic effect. Preservation of osteocyte life span can increase bone strength, which may be disproportional to the increase in bone mass. Pretreatment of osteocytes with bisphosphonates for 1h before the addition of 10-6 M dexamethasone inhibited glucocorticoid-induced apoptosis, with minimal . effective concentration between 10-9-10-8 M.
- IC ICM A61K031-00
- CC 1-10 (Pharmacology)

Section cross-reference(s): 63

IT 1406-16-2, Vitamin d 13598-36-2D, Phosphonic acid, alkylidenebisderivs. 32222-06-3, Calcitriol 40391-99-9 63132-38-7, IG 9402 63132-38-7D, IG 9402, salts 66376-36-1, Alendronate 121368-58-9, Olpadronate

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(increasing bone strength with selected bisphosphonates)

L41 ANSWER 8 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1999:738897 CAPLUS Full-text

DOCUMENT NUMBER:

132:59109

TITLE:

Prevention of osteocyte and osteoblast apoptosis by

bisphosphonates and calcitonin

AUTHOR(S): Plotkin, Lilian I.; Weinstein, Robert S.; Parfitt, A.

Michael; Roberson, Paula K.; Manolagas, Stavros C.;

Bellido, Teresita

CORPORATE SOURCE: Division of Endocrinology and Metabolism, Center for

Osteoporosis and Metabolic Bone Diseases, University of Arkansas for Medical Sciences, Little Rock, AR,

72205, USA

SOURCE: Journal of Clinical Investigation (1999), 104(10),

1363-1374

CODEN: JCINAO; ISSN: 0021-9738

PUBLISHER: American Society for Clinical Investigation

DOCUMENT TYPE: Journal LANGUAGE: English

ΔR Glucocorticoid-induced osteoporosis may be due, in part, to increased apoptosis of osteocytes and osteoblasts, and bisphosphonates (BPs) are effective in the management of this condition. We have tested the hypothesis that BPs suppress apoptosis in these cell types. Etidronate, alendronate, pamidronate, olpadronate, or amino-olpadronate (IG9402, a bisphosphonate that lacks antiresorptive activity) at 10-9 to 10-6 M prevented apoptosis of murine osteocytic MLO-Y4 cells, whether it was induced by etoposide, $TNF-\alpha$, or the synthetic glucocorticoid dexamethasone. BPs also inhibited apoptosis of primary murine osteoblastic cells isolated from calvaria. Similar antiapoptotic effects on MLO-Y4 and osteoblastic cells were seen with nanomolar concns. of the peptide hormone calcitonin. The antiapoptotic effect of BPs and calcitonin was associated with a rapid increase in the phosphorylated fraction of extracellular signal regulated kinases (ERKs) and was blocked by specific inhibitors of ERK activation. Consistent with these in vitro results, alendronate abolished the increased prevalence of apoptosis in vertebral cancellous bone osteocytes and osteoblasts that follows prednisolone administration to mice. These results suggest that the therapeutic efficacy of BPs or calcitonin in diseases such as glucocorticoidinduced osteoporosis may be due, in part, to their ability to prevent osteocyte and osteoblast apoptosis.

CC 1-12 (Pharmacology)

Section cross-reference(s): 2

IT 2809-21-4 9007-12-9, Calcitonin 13598-36-2D, Phosphonic acid,
 alkylidinebis- derivs. 40391-99-9 63132-38-7, IG 9402
 66376-36-1, Alendronate 121368-58-9, Olpadronate
 RL: BAC (Biological activity or effector, except adverse); BSU (Biological)

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(prevention of osteocyte and osteoblast apoptosis by bisphosphonates and calcitonin)

REFERENCE COUNT: 68 THERE ARE 68 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L41 ANSWER 9 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1986:430115 CAPLUS Full-text

DOCUMENT NUMBER:

105:30115

TITLE:

Treatment of collagenous tissue

INVENTOR(S): Dewanjee, Mrinal Kanti
PATENT ASSIGNEE(S): Mayo Foundation, USA
SOURCE: Eur. Pat. Appl., 32 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|--------|--------------|-----------------|----------|
| | | | , | |
| EP ·174737 | A2 | 19860319 | EP 1985-305681 | 19850809 |
| EP 174737 | A3 | 19870311 | | |
| R: BE, CH, DE, | FR, GB | , IT, LI, NL | , SE | |
| DK 8503667 | Α | 19860215 | DK 1985-3667 | 19850813 |
| AU 8546130 | A | 19860327 | AU 1985-46130 | 19850813 |
| AU 558688 | B2 | 19870205 | | |
| JP 61137825 | Α | 19860625 | JP 1985-179218 | 19850814 |
| PRIORITY APPLN. INFO.: | | | US 1984-640725 | 19840814 |

AB A process is given for the treatment of collagenous tissue to adapt it for use as a prosthetic implant and to promote the growth of endothelial cells thereon. The tissue is treated with at least 1 surfactant to remove deleterious material and open up the fibrous structure of the collagenous tissue, washed, fixed with glutaraldehyde, and the glutaraldehyde-fixed tissue is treated with a calcification-inhibiting agent, an agent that inhibits infiltration and attack by phagocytic cells and/or an agent that inhibits infection; and then the resulting matrix is treated with a reducing agent to stabilize the bonding of the agents and glutaraldehyde to the tissue. calf pericardial tissue was kept in Triton X100 for 3 h, washed, placed in 0.5% glutaraldehyde in 0.05M acetate buffer (pH 5.5) for 3.5 h, rinsed, and immersed in 3-amino-1-hydroxypropane-1,1- diphosphonic acid (16 mg/mL) in 0.05M acetate buffer, for 2-3 h. After 3 added cycles of immersion in glutaraldehyde (12 h)/rinsing, the tissue was soaked in 5 mg NaBH4/mL for 30 min, rinsed, and stored in 0.5% glutaraldehyde. When valves made of this tissue were implanted in calves, there was no calcification, and abundant endothelial cell growth was observed

IC ICM A61L027-00

ICS A01N001-02

63-7 (Pharmaceuticals) CC

151-21-3, biological studies TТ 59-05-2 61-24-5 9005-65-6 9063-89-2

63132-37-6 63132-38-7 79778-41-9 40391-99-9 97815-71-9

RL: BIOL (Biological study)

(collagenous tissue treatment with, for transplants)

L41 ANSWER 10 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1983:181876 CAPLUS Full-text

DOCUMENT NUMBER:

98:181876

TITLE:

Scale prevention with special reference to threshold

treatment

AUTHOR (S) :

Van Rosmalen, G. M.

CORPORATE SOURCE:

Dep. Chem., Delft Univ. Technol., Delft, 2628 RZ,

Neth.

SOURCE:

Chemical Engineering Communications (1983), 20(3-4),

CODEN: CEGCAK; ISSN: 0098-6445

DOCUMENT TYPE:

Journal English

LANGUAGE:

Various chemical, phys., and mech. methods for preventing deposition of mineral scale are described. The suitability of the different methods, which largely depends on the specific features and requirements of the system involved, is discussed. Special emphasis is placed upon the threshold treatment, where the growth process is retarded by the addition of trace amts. of growth inhibitors. Growth expts. were performed on BaSO4 and CaSO4.2H2O seed crystals, suspended in a supersatd. solution with and without organic bisphosphonates as inhibitors. Two methods are selected for the anal. of the growth data. A degree of inhibition is defined to obtain a quant. description of the growth-inhibitor effect on the growth rate. The effect of the mol. structure of various bisphosphonates is shown. The effect of a bisphosphonate on the geometry of CaSO4.2H2O crystals is illustrated.

CC 48-11 (Unit Operations and Processes)

Section cross-reference(s): 61

TT 2809-21-4 29712-28-5 51395-42-7 63132-38-7 63161-30-8

RL: USES (Uses)

(crystal growth rates of barium sulfate and calcium sulfate dihydrate in relation to)

L41 ANSWER 11 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN

1983:597742 CAPLUS Full-text ACCESSION NUMBER:

DOCUMENT NUMBER: 99:197742

TITLE: The influence of various phosphonates on the growth

rate of barium sulfate crystals in suspension

AUTHOR (S): Van der Leeden, M. C.; Reedijk, J.; Van Rosmalen, G.

CORPORATE SOURCE: Dep. Chem., Delft Univ. Technol., Delft, 2628 RZ,

Neth.

SOURCE: Estudios Geologicos (Madrid) (1982), 38(3-4), 279-87

CODEN: EGLMA9; ISSN: 0367-0449

DOCUMENT TYPE:

Journal

LANGUAGE: English

AB The growth of BaSO4 crystals on walls of petrochem. equipment can be slowed by phosphonates having dissociated PO32- groups and H-bonding groups, such as CO2H, OH, or/and NH3+ groups. HO2CCH2CH(CO2H)CH(PO3H2)2 [51395-42-7] and MeC(OH)(PO3H2)2 [2809-21-4] are especially effective.

CC 51-23 (Fossil Fuels, Derivatives, and Related Products)

2809-21-4 29712-28-5 IT 29712-30-9 32545-63-4 40391-99-9 51395-42-7

53818-08-9 63132-38-7 63161-30-8 73514-83-7

RL: USES (Uses)

(barium sulfate scale inhibitors, for petrochem. equipment)

L41 ANSWER 12 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1980:181274 CAPLUS Full-text

DOCUMENT NUMBER:

92:181274

TITLE:

SOURCE:

Synthesis of 2- and 3-substituted alkanediphosphonic

acids

AUTHOR(S):

Worms, K. H.; Blum, H.; Hempel, H. U.

CORPORATE SOURCE:

Henkel KGaA, Duesseldorf, D-4000/1, Fed. Rep. Ger. Zeitschrift fuer Anorganische und Allgemeine Chemie

(1979), 457, 214-18

CODEN: ZAACAB; ISSN: 0044-2313

DOCUMENT TYPE:

Journal

LANGUAGE:

German

AB Approx. 15 title compds. were prepared by phosphonylation of, primarily, aminoalkanoic acids and aminoalkanonitriles. Thus, 1 mol H3PO3, 1 mol PC13, 330 mL PhCl and 0.5 mol Et2N(CH2)2CO2H gave 57% Et2N(CH2)2C(OH)[P(O)(OH)2]2. Similarly prepared were Me2C(NH2)C(OH)[P(O)(OH)2]2, MeCH(NH2)CH2C(OH)[P(O)(OH)2]2, H2NCH2CH2C(OH)[P(O)(OH)2]2, and H2NCHPhCH2C(OH)[P(O)(OH)2]2. Phosphonylation of 0.25 mol H2NCH2CH2CN in 100 mL dioxane with 0.5 mol PBr3 followed by hydrolysis gave H2NCH2CH2C(NH2)[P(O)(OH)2]2. Similarly prepared, were MeCH(NH2)CH2C(NH2)[P(O)(OH)2]2, MeNHCH2CH2C(NH2)[P(O)(OH)2]2, and Me2NCH2CH2C(NH2)[P(O)(OH)2]2.

29-7 (Organometallic and Organometalloidal Compounds) CC

· IT 63132-38-7P

> RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reaction with nitrous acid)

L41 ANSWER 13 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1977:468491 CAPLUS Full-text DOCUMENT NUMBER:

87:68491

TITLE:

1-Hydroxy-3-aminoalkane-1,1-diphosphonic acids

INVENTOR(S):

Blum, Helmut; Worms, Karl Heinz

PATENT ASSIGNEE(S):

Henkel und Cie. G.m.b.H., Fed. Rep. Ger.

SOURCE:

Ger. Offen., 13 pp. CODEN: GWXXBX

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | | DATE |
|------------------------|------------|-----------|-----------------|---|----------|
| | | | | | |
| DE 2534391 | A1 | 19770217 | DE 1975-2534391 | | 19750801 |
| DE 2534391 | _ C2 | 19830113 | | | |
| NL 7607703 | Α | 19770203 | NL 1976-7703 | | 19760712 |
| US 4054598 | Α | 19771018. | US 1976-705792 | | 19760716 |
| BE 844649 | A1 | 19770131 | BE 1976-169348 | | 19760729 |
| JP 52019628 | Α | 19770215 | JP 1976-91196 | | 19760730 |
| JP 59025798 | В | 19840621 | | | |
| CH 599234 | A 5 | 19780531 | CH 1976-9788 | | 19760730 |
| GB 1540238 | A | 19790207 | GB 1976-31891 | | 19760730 |
| AT 349642 | В | 19790410 | AT 1976-5633 | | 19760730 |
| AT 350161 | В | 19790510 | AT 1976-5631 | | 19760730 |
| AT 7605631 | Α | 19781015 | | | |
| CH 620359 | A 5 | 19801128 | CH 1976-9789 | | 19760730 |
| PRIORITY APPLN. INFO.: | | | DE 1975-2534391 | Α | 19750801 |

AB RCH2C(OH)[P(O)(OH)2]2 (I, R = Me2NCH2, Et2NCH2, H2NCHMe) were prepared e.g. by treating RCH2CO2H with H3PO3 and P trihalide. I complex with 710 to 2500 mg CaCO3/g I at pH 11.

/ IC C07F009-38

CC 29-7 (Organometallic and Organometalloidal Compounds)

IT 63132-38-7

RL: RCT (Reactant); RACT (Reactant or reagent)
 (diazotization and hydrolysis of)

L41 ANSWER 14 OF 15 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1977:423490 CAPLUS $\underline{\text{Full-text}}$

DOCUMENT NUMBER:

87:23490

TITLE: 1,3-Diaminoalkane-1,1-diphosphonic acids

INVENTOR(S):

Blum, Helmut; Worms, Karl Heinz

PATENT ASSIGNEE(S):

Henkel und Cie. G.m.b.H., Fed. Rep. Ger.

SOURCE:

Ger. Offen., 14 pp.

CODEN: GWXXBX

DOCUMENT TYPE:

Patent German

LANGUAGE:
FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|-------------|------|----------|-----------------|----------|
| | | | | |
| DE 2534390 | A1 | 19770217 | DE 1975-2534390 | 19750801 |
| DE 2534390 | C2 | 19830113 | | |
| NL 7607702 | A | 19770203 | NL 1976-7702 | 19760712 |
| NL 186575 | В | 19900801 | | |
| NL 186575 | С | 19910102 | · | |
| BE 844648 | A1 | 19770131 | BE 1976-169347 | 19760729 |
| JP 52019627 | A | 19770215 | JP 1976-91195 | 19760730 |
| JP 59025797 | В | 19840621 | | |
| FR 2319645 | A1 | 19770225 | FR 1976-23296 | 19760730 |

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CH 599233
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                                                                    19760730
     AT 347591
                                            AT 1976-5632
                          В
                                19790110
                                                                    19760730
     GB 1540239
                          Α
                                19790207
                                            GB 1976-31892
                                                                    19760730
     AT 350160
                          В
                                19790510
                                            AT 1976~5630
                                                                    19760730
     AT 7605630
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     CH 620358
                          A5
                                19801128
                                            CH 1976-9787
                                                                    19760730
PRIORITY APPLN. INFO.:
                                            DE 1975-2534390
                                                                 A 19750801
AB
     RCH2C(NH2)[P(O)(OH)2]2 (I, R = Me2NCH2, MeNHCH2, H2NCH2, H2NCHMe) were
     prepared by treating RCH2CN with PBr3 and H2O. I complex 630->2500 mg CaCO3/g
     I at pH 11 and can be used for water softening and in the treatment of
     calcification disorders.
IC
     C07F009-38
CC
     29-7 (Organometallic and Organometalloidal Compounds)
     Section cross-reference(s): 61, 63
IT
     63132-36-5P
                   63132-37-6P 63132-38-7P
                                             63161-35-3P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation and complexing properties of)
```

```
ANSWER 15 OF 15 PHAR COPYRIGHT 2007 Informa UK Ltd on STN
L41
AN
     14466 PHAR
DN
     025567
ED
     Entered STN: 23 Apr 2003
     Last Updated on STN: 15 Dec 2003
CN
     lidadronate
CN
     IG-9402
CN
     Phosphonic acid, (1-amino-3-(dimethylamino)propylidene)bis- (CAS)
     63132-38-7
MF
     C5 H16 N2 O6 P2
MW
     262.14
HAC
HD
     6
LOGP -0.74
FRB
    9
NCE Yes
STA Ceased
```

CO

PI WO 9702827

PRAI GB 19950707

SO Pharmaprojects. PJB Publications, T&F Informa UK Ltd, London Lidadronate (IG-9402) was under development by Gador for the treatment of urolithiasis, osteoporosis, periodontal diseases and other bone disorders (Direct communication, Gador, 23 Jan 2001). It is orally-active, and inhibits glucocorticoid-induced osteocyte apoptosis (Direct communications, Gador, 25 Jan 1999 and 28 Jan 2002).

Preclinical

In vivo, it showed no interference with resorptive bone metabolism (Direct communication, Gador, 28 Jan 2002). Lidadronate and analogues are patented as selective modulators of osteoblast-osteocyte cells (Direct communication, Gador, 25 Jan 1999). Updated by JB on

19/2/2002.

DSTA World: No Development Reported Argentina: Preclinical Licensing Availability Unknown Australia: Licensing Availability Unknown Austria: Licensing Availability Unknown Belgium: Licensing Availability Unknown Brazil: Licensing Availability Unknown Canada: Licensing Availability Unknown Chile: Licensing Availability Unknown China: Licensing Availability Unknown Colombia: Licensing Availability Unknown Denmark: Licensing Availability Unknown Finland: Licensing Availability Unknown France: Licensing Availability Unknown Germany: Licensing Availability Unknown Greece: Licensing Availability Unknown Hong Kong: Licensing Availability Unknown India: Licensing Availability Unknown Ireland: Licensing Availability Unknown Israel: Licensing Availability Unknown Italy: Licensing Availability Unknown Japan: Licensing Availability Unknown Luxembourg: Licensing Availability Unknown Malaysia: Licensing Availability Unknown Mexico: Licensing Availability Unknown Netherlands: Licensing Availability Unknown New Zealand: Licensing Availability Unknown Norway: Licensing Availability Unknown Peru: Licensing Availability Unknown Philippines: Licensing Availability Unknown Portugal: Licensing Availability Unknown Russian Federation: Licensing Availability Unknown South Africa: Licensing Availability Unknown Korea, Republic of: Licensing Availability Unknown Spain: Licensing Availability Unknown Sweden: Licensing Availability Unknown Switzerland: Licensing Availability Unknown Thailand: Licensing Availability Unknown Turkey: Licensing Availability Unknown United Kingdom: Licensing Availability Unknown United States: Licensing Availability Unknown Venezuela: Licensing Availability Unknown CC G4Z Urological A1A Stomatological Osteoporosis treatment CTIndication: Unspecified (No Development Reported) GEN Target Gene: Unspecified ORGM CH-SY (Chemical, synthetic) RTE A-PO (Alimentary, po) RDAT 20031211 RNTE ##Actual; No Development Reported 20010123 ##Estimated; Names Granted IG-9402 19970415 ##Estimated; New Product in Pharmaprojects PHCD OSTEOBL-AG; Osteoblast stimulant; Physiological, Biochemical, Osteoblast stimulant; P-B-OSTEOBL-AG. PHCD OSTEOBL-AN; Osteoblast inhibitor; P=Biochemical, OSTEO; P=B-OSTEOCL-AN. PHCD APOP-AN; Apoptosis antagonist; Physiological, Biochemical, Apoptosis antagonist; General apoptosis antagonist; Apoptosis inhibitor; P-B-APOP-AN. PHCD P; P-B; P-B-OSTEOBL; P-B-OSTEOBL-AG; P-OSTEOBL; P-OSTEOBL-AG; P-AG;

B; B-OSTEOBL; B-OSTEOBL-AG; B-AG; OSTEOBL; OSTEOBL-AG; P-B-AG; P=B; P=B-OSTEOCL; P=B-OSTEOCL-AN; P=B-AN; OSTEOCL; OSTEOCL-AN; P-B-APOP; P-B-APOP-AN; P-APOP; P-APOP-AN; P-APOP, B-APOP-AN; B-APOP; APOP-AN; P-B-AN.

| | Pharmacology (PHCD) | Status | (DSTC) |
|-----|-------------------------------|--------|--------|
| G4Z | OSTEOBL-AG OSTEOBL-AN APOP-AN | | |
| A1A | OSTEOBL-AG OSTEOBL-AN APOP-AN | N | |
| M5A | OSTEOBL-AG OSTEOBL-AN APOP-AN | N | |

LCDAT 20031211: IL : No development reported

FILE 'HOME' ENTERED AT 15:33:00 ON 05 SEP 2007

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=> d his nofile
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(FILE 'HOME' ENTERED AT 15:15:44 ON 05 SEP 2007)
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FILE 'CAPLUS' ENTERED AT 15:16:08 ON 05 SEP 2007

E US2003-619729/APPS

L1 1 SEA ABB=ON US2003-619729/AP

D SCAN

FILE 'REGISTRY' ENTERED AT 15:16:56 ON 05 SEP 2007

L2 STR

L6

L13

L15

L3 0 SEA SSS SAM L2

FILE 'CAPLUS' ENTERED AT 15:19:07 ON 05 SEP 2007

SEL RN L1

L4 220 SEA ABB=ON ROLDAN E?/AU

L5 2651 SEA ABB=ON PEREZ LLORET A?/AU OR PEREZ A?/AU OR LLORET A?/AU

402 SEA ABB=ON VAZQUEZ G?/AU

L7 198 SEA ABB=ON BOLAND R?/AU

L8 138 SEA ABB=ON PAPAPOULOS S?/AU

L9 1 SEA ABB=ON L4 AND L5 AND L6 AND L7 AND L8

D SCAN

FILE 'REGISTRY' ENTERED AT 15:20:27 ON 05 SEP 2007

L10 21 SEA ABB=ON (63132-38-7/BI OR 121368-58-9/BI OR 13598-36-2/BI OR 1406-16-2/BI OR 19356-17-3/BI OR 21829-25-4/BI OR 27121-73-9

/BI OR 2809-21-4/BI OR 32222-06-3/BI OR 40391-99-9/BI OR 471-34-1/BI OR 50-14-6/BI OR 52-53-9/BI OR 67-97-0/BI OR

7440-70-2/BI OR 7693-13-2/BI OR 7782-41-4/BI OR 84449-90-1/BI

OR 9001-86-9/BI OR 9002-64-6/BI OR 9002-72-6/BI)

D SCAN

E "PHOSPHONIC ACID, P,P'-(1-AMINO-3-(DIMETHYLAMINO)PROPYLIDENE)

L11 1 SEA ABB=ON "PHOSPHONIC ACID, P,P'-(1-AMINO-3-(DIMETHYLAMINO)PR

OPYLIDENE) BIS-"/CN

D RN

E 63132-38-7/CRN

L12 2 SEA ABB=ON 63132-38-7/CRN

3 SEA ABB=ON (L11 OR L12)

D SCAN

FILE 'ZCAPLUS' ENTERED AT 15:23:17 ON 05 SEP 2007

L14. 24 SEA ABB=ON L13

FILE 'REGISTRY' ENTERED AT 15:23:27 ON 05 SEP 2007

STR 63132-38-7

L16 0 SEA FAM SAM L15

L17 21 SEA FAM FUL L15 EXTEND

L18 3 SEA FAM FUL L15

SAVE TEMP L18 ISS729FAM/A

L19 3 SEA ABB=ON L18 AND L13

D LC 1-3

FILE 'REGISTRY' ENTERED AT 15:25:19 ON 05 SEP 2007

SET TERMSET E#

DEL SEL Y

SEL L19 3 RN

L20 1 SEA ABB=ON 63132-38-7/RN

SET TERMSET LOGIN

```
FILE 'PHAR' ENTERED AT 15:25:23 ON 05 SEP 2007
     1 SEA ABB=ON L20
L21
                SET LINE 250
                SET DETAIL OFF
                SET LINE LOGIN
                SET DETAIL LOGIN
                D SCAN
                D TRIAL
     FILE 'STNGUIDE' ENTERED AT 15:25:52 ON 05 SEP 2007
     FILE 'CAPLUS' ENTERED AT 15:27:36 ON 05 SEP 2007
L22
           24 SEA ABB=ON L13
L23
            10 SEA ABB=ON (L4 OR L5 OR L6 OR L7 OR L8 OR L1) AND L22
   FILE 'BIOSIS' ENTERED AT 15:28:13 ON 05 SEP 2007
           264 SEA ABB=ON ROLDAN E?/AU
L24
         2810 SEA ABB=ON PEREZ LLORET A?/AU OR PEREZ A?/AU OR LLORET A?/AU 332 SEA ABB=ON VAZQUEZ G?/AU
L25
L26
L27
           231 SEA ABB=ON BOLAND R?/AU
L28
           278 SEA ABB=ON PAPAPOULOS S?/AU
L29
             2 SEA ABB=ON L13
             2 SEA ABB=ON (L24 OR L25 OR L26 OR L27 OR L28) AND L29
L30
   FILE 'DRUGU' ENTERED AT 15:29:06 ON 05 SEP 2007
           15 SEA ABB=ON ROLDAN E?/AU
L31
           224 SEA ABB=ON PEREZ LLORET A?/AU OR PEREZ A?/AU OR LLORET A?/AU
L32
L33
            23 SEA ABB=ON VAZQUEZ G?/AU
            22 SEA ABB=ON BOLAND R?/AU
L34
            43 SEA ABB=ON PAPAPOULOS S?/AU
L35
L36
            1 SEA ABB=ON L13
               D TRIAL
L37
             O SEA ABB=ON L36 AND LITERATURE/FS
     FILE 'CAPLUS' ENTERED AT 15:31:22 ON 05 SEP 2007
               D QUE NOS L23
     FILE 'BIOSIS' ENTERED AT 15:31:23 ON 05 SEP 2007
               D QUE NOS L30
     FILE 'CAPLUS, BIOSIS' ENTERED AT 15:31:27 ON 05 SEP 2007
            11 DUP REM L23 L30 (1 DUPLICATE REMOVED)
L38
                    ANSWERS '1-10' FROM FILE CAPLUS
                    ANSWER '11' FROM FILE BIOSIS
               D IBIB ABS HITSTR 1-11
     FILE 'REGISTRY' ENTERED AT 15:32:00 ON 05 SEP 2007
               D STAT QUE L13
               D IDE L13 1-3
     FILE 'CAPLUS' ENTERED AT 15:32:32 ON 05 SEP 2007
              D QUE L22
L39
           14 SEA ABB=ON L22 NOT L23
     FILE 'BIOSIS' ENTERED AT 15:32:32 ON 05 SEP 2007
               D QUE L29
L40
              0 SEA ABB=ON L29 NOT L30
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FILE 'PHAR' ENTERED AT 15:32:32 ON 05 SEP 2007

D QUE L21

FILE 'CAPLUS, PHAR' ENTERED AT 15:32:38 ON 05 SEP 2007
L41

15 DUP REM L39 L21 (0 DUPLICATES REMOVED)

ANSWERS '1-14' FROM FILE CAPLUS

ANSWER '15' FROM FILE PHAR

D IBIB ABS HITIND 1-14

D IBIB ABS HITIND 1-14 D ALL 15

FILE 'HOME' ENTERED AT 15:33:00 ON 05 SEP 2007